AMENDMENTS TO THE SPECIFICATION

Please enter the following amendments to the specification. For the Examiner's convenience, the following amendments include markings to show the changes to a portion of the specification relative to the immediate prior version of that portion of the specification.

REPLACE Paragraph [0005] with the Following:

[0005] Using data describing contraindications and complications from FDA guidelines, public manufacturer data, clinical results from professional industry conferences, interdisciplinary scientific literature and negative case law, as well as dynamic data about surgeons' outcome and results history for the surgery, the method calculates in real time a customized risk assessment in terms of the probability of a successful outcome for the patient undergoing a surgery or treatment by a certain physician using a selected device devices or therapy. Based on the customized risk analysis, the method causes a consent form to be generated in real time, which comprises standardized and individualized paragraphs explaining the risks associated with the surgery or treatment for the patient.

REPLACE Paragraph [00016] with the Following:

[00016] Fig. 1 illustrates a prior art method for providing a patient an informed consent form. As shown, the prior process includes information on the absolute contraindications 10 of refractive laser surgery for correcting vision, which is gathered from the medical literature, FDA guidelines, the manufacturer's warnings and professional conferences. These absolute contraindications 10 have traditionally been used to formulate informed consent forms and to assess risks for patients considering the procedure at the point 100 when the surgery commences on the general public.

REPLACE Paragraph [00017] with the Following:

[00017] As Fig. 1 shows, once the "informed" consent is formulated, there is little or no opportunity in the prior art process to re-formulate the informed consent so as to include the mounting evidence 30 of complications resulting from the procedure. That is, the basis for the informed consent form in the prior art method constitutes almost entirely the absolute contraindications 10 originally formulated before actual practice of the surgical technique on the public.

REPLACE Paragraph [00019] with the Following:

[00019] Fig. 2 illustrates an exemplary embodiment of the present method in which the data is continually accumulated, processed, and presented to the patient to provide an up to the moment truly informed consent. A contracted health care provider 280, such as a hospital, a clinic, physician's practice group or a sole practitioner contracts to become a participating member. Participation in the present process provides members and their patients the ability to calculate and print out an individualized risk assessment for undergoing surgery. Semi-static data 200, which comprises information on absolute contraindications 210 and emerging contraindications 220, emerging data relating to complications 230 occurring as a result of the surgery, and unknown contraindications 240 are input into the rule-based, informed consent engine 250. The method then calculates, using a clinical basis process 260 (shown in more detail in Figure 3), a customized risk analysis for the patient contemplating a procedure.

REPLACE Paragraph [00021] with the Following:

[00021] Fig. 3 illustrates an embodiment of the clinical basis process 260 [[300]] in which Semi-static data 200 and Dynamic data 350 are used to generate the rule-based algorithm 390, which shapes the <u>customized</u> Dynamic Informed Consent Form 270. The algorithm 390 1.111 Amendment dated May 2, 2007

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formulates rules of risks relating to the surgical procedure 380 and rules of risks developed from analyzing data on post-operative events and outcomes 370. In turn, the generated rules 370 and 380 do not remain static but are re-evaluated and re-generated upon the input of data 310 on patients' pre-operative care, data 340 on patients' [[and]] post-operative care [[340]], data 320 on the surgical procedure, and data 330 on the positive and negative outcomes of the surgery as performed by surgeons associated with the health care provider.

REPLACE Paragraph [00022] with the Following:

[00022] The clinical basis process 260 [[300]] is a real time, iterative calculation of the risks for an individual patient, considering not only the semi-static data 200 of contraindications but also the dynamic data 350 on complications and emerging contraindications, which relate to the outcome and result track record of particular surgeons associated with a contracted health care provider. In other words, the clinical basis process 260 [[300]] evaluates an individualized patient risk assessment from all the dynamic data on patient outcomes and results of surgeries performed by various surgeons of a contracted provider 280 as well as from the semi-static data 200 relating to surgical devices used for the procedure. The same process is equally adapted to the evaluation and risk assessment related to other medical therapeutic processes and treatments and those who provide them.

REPLACE Paragraph [00023] with the Following:

Once calculated, the risk assessment is presented in the form of a customized dynamic informed consent form 270. This is accomplished by drafting separate sentences or paragraphs that explain different risks and creating a calculus that associates different explanatory sentences or paragraphs with different patient conditions and for different surgeons. 1.111 Amendment dated May 2, 2007 Reply to Office Action dated January 3, 2007

REPLACE Paragraph [00024] with the Following:

[00024] The method is iterative and dynamic in that the clinical basis process 260 [[300]] may be continually updated with real time data to provide a continually updated rule-based algorithm 390. That is, as updated data on surgeons' and patients' outcomes and results history are acquired, the rule-based algorithm 390 both fine-tunes the surgical risks associated with various patient conditions and different surgeons and updates the calculus that associates the risk explanations with these conditions and surgeons. By updating the risk assessment algorithm and the calculus for associating explanations with surgery conditions particularly by inputting dynamic data 350, the resultant informed consent form may be continually updated and customized for individual patients.

REPLACE Paragraph [00026] with the Following:

[00026] An algorithm-based data engine dynamically processes and analyzes static and dynamic data on a continuing, recursive basis. The result is a presentation in real-time that identifies risk in surgical and medical procedures that provides a patient with a live, state of the moment informed consent based on existing indications and warnings, literature, and iteratively processed data gathered from other patients. The process can also generate warnings based on all available data when such warnings would be appropriate.